PATENT SPECIFICATION

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C3Q C13A C4A D1A T2A (72) Inventor VISVALDIS ABOLINS and FRED FRANK HOLUB

(54) REINFORCED THERMOPLASTIC COMPOSITIONS

(71) We, GENERAL ELECTRIC COMPANY, a corporation organised and existing under the Laws of the State of New York, United States of America, of 1 River Road, Schenectady 12305, New York, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

This invention relates to reinforced thermoplastic compositions for molding and conversion to films, fibers, and sheets. Also contemplated are flame retardant

and non-dripping forms of such compositions.

High molecular weight linear polyesters and copolyesters of glycols and terephthalic or isophthalic acid have been available for a number of years. These are described *inter alia* in Whinfield *et al.*, U.S. Patent No. 2,465,319 and in Pengilly, U.S. Patent No. 3,047,539. These patents disclose that the polyesters are particularly advantageous as film and fiber formers.

The most commonly employed polyester prepared by these teachings, poly(ethylene terephthalate), has only fairly recently been widely accepted for use as a molding resin because of its relative brittleness in thick sections when crystallized from the melt. The problem was overcome by varying the crystal texture, e.g., by using two-step molding cycles or by including nucleating agents, and by molecular weight control. This permitted the marketing of injection-moldable poly(ethylene terephthalates) which typically, in comparison with other thermoplastics, offer a high degree of surface hardness and abrasion resistance, and lower surface friction. Solvent resistance is outstanding, making such compositions uniquely useful in the automotive and signaff industries. Among the

compositions uniquely useful in the automotive and aircraft industries. Among the disadvantages of poly(ethylene terephthalate) in molding compositions are the relatively high cost of manufacture (because of the need to control crystal texture), a relatively low degree of moisture resistance, a rapid burning rate and a tendency to drip flaming resin while burning.

Surprisingly, it has been found that the need to employ nucleating agents or

two stage molding cycles to avoid brittleness in thick molded sections is obviated if higher homologs of poly(ethylene terephthalate) are mixed with it or substituted for it. For example, within the broad disclosure of the Whinfield et al and Pingilly patents there exists a family of polyester resins which are normally crystalline and which rapidly crystallize from the melt. These crystallize so rapidly, in fact, that standard injection molding cycles can be used and there is no need to include nucleating agents in the formulation. Parts molded from such polyesters, in contrast to those of poly(ethylene terephthalate) alone, have good impact strength

The rapidly crystallizable higher homologs comprise generally polyester resins which are poly(alkylene terephthalates, isophthalates or mixed terephthalates and isophthalates), wherein the alkylene groups contain from 3 to 10, and especially 3 to 6, carbon atoms.

Simultaneously with the development of injection molding grades of



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	poly(ethylene terephthalate) resins, fiber glass reinforced compositions were also provided. See Furukawa et al., U.S. Patent No. 3,368,995. These injection moldable compositions provided all of the advantages of the file.	
_	moldable compositions provided all of the advantages of unfilled polyesters and, also, because of the glass reinforcement, the molded articles had higher rigidity, yield strength, modulus and impact strength.	
5	yield strength, modulus and impact etrangely included articles had higher rigidity,	
	110WEVEL DECAUSE they used notwethylang to an habitation in the	5
	compositions required two stage molding or the use of nucleating agents to avoid embrittled thickly molded sections. Moreover, the use of nucleating agents to avoid	
	embrittled thickly molded sections. Moreover, they retained the disadvantages inherent in polyesters generally having a relatively likely and the disadvantages	
10	inherent in polyesters generally, having a relatively high cost of manufacture and formulation, and low degree of moisture resistance that of manufacture and	
	formulation, and low degree of moisture resistance, high flammability and a tendency to drip while burning.	10
	Although substitution of the smills and the	•••
	poly(ethylene terephthalate) would be expected to overcome the need for two- stage molding cycles or nucleating agents such corrections the need for two-	
15	stage molding cycles or nucleating agents, such compositions, in common with	
1.0	those described by Furukawa et al, would still be expensive, unstable to moisture, flammable and would drip while burning. While registers to a unstable to moisture,	15
	flammable and would drip while burning. While resistant to gasoline, jet fuels, and hydraulic fluids, such compositions would be of limited utilization, jet fuels, and	15
	hydraulic fluids, such compositions would be of limited utility in the automotive and aircraft industries because of their flammability.	
	of moisture sensitivity, their use in his heart family. On the other hand, because	
20	This invention relates to a thermolist, laundries or snips would be limited.	
	crystallizable polyester (as hereinafter defined) with at least one other polymer and a reinforcing filler. Without the filler component great difficulty in the polymer and	20
	a reinforcing filler. Without the filler component, great difficulty is experienced in processing the blend of two polymers. Surprisingly, as little and the second of two polymers.	
	processing the blend of two polymers. Surprisingly, as little as 2° by weight of reinforcing filler, such as glass provides uneverted in the control of the polymers.	
25	bility, surface appearance and physical processed improvement in processa-	
		25
	terephthalate) will be enhanced according to this invention by forming a reinforced combination with polyolefins and the strength rigidity and forming a	
	reinforced combination with polyolefins and the strength, rigidity, and resistance to heat distortion of polyolefins will be improved by forming a	
30	to heat distortion of polyolefins will be improved by forming a reinforced combination with poly (1,4-butylene terephthalate).	
	It is surprising and unexpected to be a late.	30
	combinations with a remarkably broad range of polymers. Usually, when such mixtures, including at least one highly crystallizable networks.	
	mixtures, including at least one highly crystallizable polymer, are molded and cooled, they are non-uniform and tend to delawiness.	
35	cooled, they are non-uniform and tend to delating polymer, are molded and individual components). In contrast to the averaged by the separate into their	
	able polyesters have been found to the expected denaviour, rapidly crystalliz-	35
	amorphous and partially crystalline, and partially composites with crystalline,	33
	reinforcement, the resulting composites are easily moldable and extrudable into	
10	articles which are uniform and which do not delaminate. All of these characteristics of the new compositions of this invention indicate that	
N.	istics of the new compositions of this invention indicate that an intercrystalline (i.e.	40
	homogeneous crystalline mixture) is formed between that an intercrystalline (i.e. co-blended polymer. In addition, the enhancement is at a little polymer and the second	40
	evidence that the reinforcing filler a mantenent in strength of the composite is	
_	silicate, titanate, carbon black, clay or glass, is unexpectedly and strongly bonded into the combination.	
5	into the combination.	
	This invention provides a reinforced thermoplastic composition that is rigid at temperatures of up to 90°F, which comprises a homeon that is rigid at	45
	rigid, reinforceable mixture of	
	(a) at least one high molecular maintains	
0	which rapidly crystallizes from the melt (as hereinbefore defined); (b) at least one high molecular weight hereinbefore defined);	
	(b) at least one high molecular weight normally crystalline, normally partially crystalline and normally crystalline, normally	50
	amorphous or normally partially crystalline amorphous polymer of: (i) at least one alignatic ethylerically amorphous polymer of:	
	propylene: unsaturated monomer excluding	
5	(ii) at least one difunctionally reactive company to the	
	self-condensation; and/or	55
	(iii) at least two difunctionally reactive compounds which are polymerizble by polycondensation;	
	in the concentration range of from 1 as on	
)	in the concentration range of from 1 to 99 parts by weight of (a) to 99 to 1 parts by weight of (b); and	
	(c) from 2 to 90° by weight based on south	60
	for said homogeneous, crystalline, normally rigid, reinforceable mixture;	
	and when polymer (b) is a polyester, it is a wholly aromatic polyester and different from polyester (a).	
i	With respect to the polyecter and	
	With respect to the polyester resin component (a), there will be selected a	65

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high molecular weight normally crystalline polycondensation product of a difunctional organic alcohol or reactive derivative thereof and a difunctional organic acid or a reactive derivative thereof. The difunctional alcohol and the difunctional acid can be wholly aliphatic or wholly aromatic or partially aliphatic and partially aromatic in character, the organic groups being straight chained, or branched, cyclic or polycyclic and containing from 2 to 20, preferably from 3 to 10, especially preferably from 3 to 6, carbon atoms. The organic groups can be unsubstituted or substituted with conventional substituents, such as alkyl, halogen, carboxyl, nitro, cyano, amido or imido groups.

While a wide variety of polyesters are suitable for this invention, it is important that they be normally crystalline, high enough in molecular weight to form films and fibers (although this is merely a measure of a suitable molecular weight and not a limitation on ultimate use) and, most importantly, rapidly crystallizable from the melt.

Polyesters which meet the criterion of being rapidly crystallizable from the melt will be those which can be injection molded into a mould having a temperature of from 160 to 175°F, preferably 170°F, when they themselves have a higher temperature than the mould in a standard, short cycle time and produce a rigid workpiece which is highly crystalline throughout. As is described in Furukawa et al. U.S. Patent No. 3,368,995, poly(ethylene terephthalate) will not produce such a workpiece because of its high crystalline melting point and low heat conductivity. Because such a polyester resin is not rapidly crystallizable, the outer part of the workpiece is amorphous, and the inner part, which is gradually cooled, is crystalline.

One convenient way to determine if a polyester is suitable for use in this invention is to injection mold it with the mold temperature of 160-175°F. in a standard short cycle, e.g., 10-90 seconds, into a workpiece about 1 inch thick. If the moulded article is rigid on removal from the mould and the inner and outer parts of the piece are uniformly milky or white and opaque and if the hardness, measured, e.g., with a Rockwell M tester, is uniform throughout, crystallization from the melt is rapid enough to be suitable. If, on the other hand, the workpiece is clear, transparent or semi-transparent (i.e., amorphous) on the outside and milky, white or opaque (i.e., crystalline) only on the inside; if the hardness is non-uniform throughout; if the workpiece is relatively soft on removal from the mould and brittle after cooling; or if uniformity can be achieved only by using a two stage molding cycle, or higher mold temperatures, e.g., 250—330°F. and long times, e.g., 90—400 sec., or by including a nucleating agent, e.g. carbon powders, metal salts or clays, then the polyester resin is not suitable for this invention.

Typical of the high molecular weight rapidly crystallizable polyester resins suitable for this invention are poly(alkylene terephthalates, isophthalates).

or mixed terephthalates and isophthalates), wherein the alkylene groups contain from 3 to 10 carbon atoms. They are available commercially or can be prepared by known techniques, such as by the alcoholysis of esters of the phthalic acid with a glycol and subsequent polymerization, by heating glycols with the free acids or with halide derivatives thereof, and similar processes. These are described in U.S. Patent No. 2,465,319 and U.S. Patent No. 3,047,539, and

Although the glycol portion of such typical polyesters can contain from 3 to 10 carbon atoms, e.g., 1,3-propylene, 1,4-butylene, 1,3-butylene, 1,2-propylene, 1,2-butylene, or 2,3-butylene, it is preferred that it contain 3 or 4 carbon atoms, in the form of linear mehylene chains.

Preferred polyesters are high molecular weight, polymeric 1,4-butylene glycol terephthalates or isophthalates having repeating units of the general formula

55 and mixtures of such esters, including copolyesters of terephthalic and isophthalic acids over the entire composition range.

Especially preferred polyesters are poly(1,4-butylene isophthalate) and poly-(1,4-butylene terephthalate). Special mention is made of the latter because it is easy to prepare from readily available materials and crystallizes at an especially rapid rate.

	1,707,273	4
5	Illustratively, sufficiently high molecular weight polyesters of the preferred type will have an intrinsic viscosity of at least 0.2 and preferably 0.4 deciliters/gram as measured in o-chlorophenol, a 60/40 phenol-tetrachloroethane mixture or a similar solvent at 25—30°C. The upper limit is not critical, but will generally be 1.5 from 0.5 to 1.3. With respect to the	5
10	With respect to the co-blending polymer (b), these will be selected from one or more of high molecular weight normally crystalline, normally amorphous or more fully below) which form with polyester component (a) a reinforceable combination with the concentration range of from 1 to 99 parts by weight of (a) to	10
15	It is to be understood that not all of the suitable polymers falling within the definition given for component (b) will form reinforceable combinations over the combinations within only portions of this range, and all non-reinforceable combinations of polyesters and such polymers within the stated range are	15
20	which persons skilled in the art will readily recognize as failing to meet the criteria set out above, i.e., they may or may not combine each other when melted, but, The co-blending polymers (b) will form an intercrystalline normally rigid	20
25	which arises from a state of molecular structure which denotes compactness of the molecular chains forming the polymer. It is attributable to the formation of solid crystals having a definite geometric for	25
30	weight polymer, depending often on the way it is produced, can exist in wholly form. Merely by way of illustration, polymers of the same alpha olefin, e.g., poly(1, but not).	30
35	butene), have different steric structures, and depending on their steric configurations, they may be crystallizable or amorphous, or mixed. Although as will be seen, polymer (b) can be either normally solid (i.e. rigid) or normally rubbery (i.e. non-rigid), the compositions of this invention are all that which ensures that the final composition will be rigid at temperatures of 75—90°F.	35
40	The polymer component (b) -file :	
45	butadiene, e.g., ethylene or an olefin of from 4 to 10 carbon atoms, e.g., butene-1, halide or vinylidene halide, for example, vinyl chloride, vinylidene chloride, or exters, alkyl methacrylates, acrylic monomers such as acrylic acid, acrylic chlorinated polyethylates. (The acrylic nitriles, such as methacrylonitrile, and	40 45
50	(ii) at least one diffunctionally reactive compound which is polymerizable by self-condensation, such as formaldehyde, trioxymethylene, phenols, e.g., 2,6- (iii) at least two diffunctionally reactive compound which is polymerizable by dimethylphenol, siloxanes, lactams, e.g. caprolactam, or aromatic sulfides; and/or	50
55	dicarboxylic acid, e.g., adipic acid, a polyorganosiloxane and an aromatic ester, a sulfone.	55
60	of (i) and (ii), or a styrene resin and (ii) such as a blended composition of polyphenylene ether; polyethylene or polybutene-I and a polyphenylene ether; or polystyrene with a wholly aromatic polyester, e.g. from	60
65	Among the preferred features of this invention are compositions where a polymer (b) is a polymerization product of at least one aliphatic ethylenically unsaturated monomer and is selected from polyethylene and other polyolefins and copolymers of such monomers, for example, polyethylene poly(methylene),	65

	normally solid count	5
5	normally solid copolymers of ethylene and butene-1, copolymers of ethylene and ethyl acrylate, or vinyl acetate, butadiene-acrylonitrile copolymers, ionomers, poly(methyl methacrylate), polyisobutylene rubbers poly(vinyl chloride), polyisobutylene rubbers poly(vinyl chloride), rubber, a rubbery copolymer of vinyl chloride with vinyl acetate, natural of butadiene and acrylonitrile. All such polymers are commercially available or can be prepared by techniques well known to those skilled in the	5
10	copolymers and terpolymers, the proportions of the repeating units may vary rubbery, or normally solid. In addition to the polymers illustrated above, other include derivatives thereof such as a liphatic ethylenically unsaturated monomers.	10
15	on trioxane, polyphenylene ethers, such as poly(2,6-dimethyl-1,4-phenylene)	15
20	of hexamethylenediamine and adipic acid, polyimides, e.g., the product maleimido diphenyl methane and methylene dianiline, normally solid or normally of the two, and conclumers of polyaryalkyl or aryl-siloxanes, or combinations	20
25	aromatic carbonates or the reaction products of bisphenol-A and iso or tere- phthaloyl chloride, as well as siloxane-nitrogen copolymers containing amido, amideimido, and imide groups. All such polymers are either commercially	25
. 30	of classes (i), (ii) and/or (iii). For example, one such mixture would comprise a high molecular weight composition which is a mixture of polystyrene or other styrene of 2,6-dimethylphenol, i.e. polysty of dimethylphenol, i.e. polystyrene of 2,6-dimethylphenol, i.e. polystyrene of 2,6-	30
35	in which the combination contains from 1 to 99 parts by weight, and preferably mixed terephthalate and isophthalate), having from 3 to 10 carbon atoms in the	35
40	either a high molecular weight poly(methyl methacrylate); a polyethylene, phenylene ether resin or a polyphenylene ether resin in combination with a styrene crystalline wholly aromatic polyester. The preferred polyester (a) is poly butylene terephthalate).	40
45	All of the present combinations include, as an essential ingredient, 2 to 90% by weight, based on total composition of a reinforcing filler (c). In general, any reinforcement can be used, e.g., fibers, whiskers or platelets of metals, e.g. aluminum, iron or nickel, and non-metals, e.g., carbon filaments, silicates.	45
50	that, unless the filler adds to the strength, stiffness and impact strength of the composition, it is only a filler and not a reinforcing filler as contemplated herein. In particular, the preferred reinforcing fillers are of glass and it is preferred to	50
55	where electrical properties are not so important, e.g., the low soda glass known as blowing, flame blowing and mechanical pulling. The preferred filaments for	55
60	The length of the glass filaments and whether or not they are bundled into given and the fibers bundled in turn to yarns, ropes or rovings, or woven into mats, compositions, it is convenient to use the filamentous glass in the form of chopped strands of from 1° to 1° long. In matical to the present strands of from 1° to 1° long. In matical to the present strands of from 1° to 1° long. In matical to the present into the present strands of from 1° to 1° long. In matical to the present into the present invention.	60
65	strands of from 1" to 1" long. In articles molded from the compositions, on the other hand, even shorter lengths will be encountered because, during compounding, considerable fragmentation will occur. This is desirable, however,	65

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	because the best properties are exhibited by thermoplastic injection molded articles in which the filament lengths lie between 0.000005" and 0.125 (1)". Preferably the sized filamentous glass reinforcement comprises from 5 to 50° by weight based on the combined weight of glass and 1.125 (1)".	
5	preferably the glass will comprise from 10 to 40% by weight based on the combined weight of glass and resin. Generally, for direct molding use, up to 60% of glass can be present without causing flow problems. However, it is useful also to prepare the	5
10	resins that are not glass reinforced to provide any desired glass content of a lower value.	10
	Because it has been found that certain commonly used flammable sizings on the glass, e.g., dextrinized starch or synthetic polymers, contribute flammability often in greater proportion than expected from the amount of the second starts.	
15	to use lightly sized or unsized glass reinforcements in those compositions of the present invention which are flame retardant. Sizings, if present, can readily be art.	15
20	It is a preferred feature of this invention also to provide flame retardant glass reinforced thermoplastic compositions, as defined above, wherein the polyester is normally flammable, the composition also including (d) a flame retardant additive in a minor proportion but in an amount at least sufficient to render the polyester resin pon-hypping or self-neighbor.	20
25	A preferred feature of the invention is a flame retardant composition as above	
	(e) a polytetrafluoroethylene resin in a minor proportion based on the composition but in an amount at least sufficient to render said polyester resin non-	25
30	When used herein the terms "non-burning", "self-extinguishing", and "non-dripping" are used to describe composites which meet the standards of ASTM test method D-635 and of the following test, hereinafter designated the SE (self-extinction test). A molded piece of about 2½" x ½" x ½" is formed from the composition. When this sample is supported vertically and ignited, if it does not form flaming droplets sufficient to ignite a piece of cotton held 12 inches beneath and extinguishes itself within an average of 5 second and	30
35	ignitions, the composition is given an SE rating of A. If the flame is extinguished within an average of 25 seconds but no greater than 30 seconds and no flaming droplets are formed, the material is given an SE rating of B. If the flame is	35
.0 .5	material is given an SE rating of C. ASTM test D-635 for flammability comprises supplied" with a Bunsen burner flame for 25 seconds, and repeating if there is no after the flame is removed, it is classified as the limit of the specimen does ignite but does not continue burning to the 4" mark	40
3	The Oxygen Index of LOI (Limiting Oxygen Index) is a direct measure of a product's combustibility, based on the oxygen content of the combustion oxygen is reduced stepwise until the material no longer supports a flame. The LOI is defined as follows:—	45
0	$LOI = \frac{100 \text{ x percentage oxygen}}{100 \text{ model}}$	
	percentage nitrogen + percentage oxygen in combustion-supporting	50

The compositions of this invention which contain flame-retardant additives in the specified amounts have a substantially higher oxygen index and thus are much less combustible than the controls.

The flame-retardant additives (d) which are used according to this invention are materials which are well known to those skilled in the art. Generally speaking, the more important of these compounds contain chemical elements employed for their ability to impart flame resistance, e.g., bromine, chlorine, antimony, phosphorus and nitrogen. It is preferred that the flame-retardant additive

comprise a halogenated organic compound (brominated or chlorinated); a halogen-containing organic compound in admixture with the antimony trioxide; elemental phosphorus (which in spite of its own flammability imparts flame-retardant properties to the compositions of this invention) or a phosphorus compound; a halogen-containing compound in admixture with a phosphorus compound or compounds containing phosphorus-nitrogen bonds or a mixture of two or more of the foregoing.

The amount of flame-retardant additive used is not critical to the invention, so long as it is present in a minor proportion based on said composition—major proportions will detract from physical properties—but at least sufficient to render the polyester resin-blend non-burning or self-extinguishing. Those skilled in the art are well aware that the amount will vary with the nature of the polymers in the blend and with the efficiency of the additive. In general, however, the amount of additive will be from 0.5 to 50 parts by weight per hundred parts of components (a) range will be from 8 to 12 parts of additive per 100 parts of (a) + (b). Smaller amounts of compounds highly concentrated in the elements responsible for flame-retardance will be sufficient, e.g., elemental red phosphorus will be preferred at form of triphenyl phosphate will be used at 25 parts of phosphate per hundred parts of (a) + (b). Halogenated aromatics will be used at 2 to 20 parts and parts of components (a) plus (b).

Among the useful halogen-containing compounds are those of the formula

wherein R is an alkylene, alkylidene or cycloaliphatic linkage, e.g., methylene, ethylene, propylene, isopropylene, isopropylidene, butylene, isobutylene, amylene, cyclohexylene, or cyclopentylidene; a linkage selected from ether; carbonyl; amine; a sulfur-containing linkage, e.g., sulfide, sulfoxide, or sulfone; carbonate; or a phosphorus-containing linkage. R can also consist of two or more alkylene or alkylidene linkages connected by such groups as aromatic, amino, ether, ester, carbonyl, sulfide, sulfoxide, sulfone, or a phosphorus-containing linkage. R can be a dihydric phenol carbonate linkage, e.g., bisphenol—A

Ar and Ar' are mono- or polycarbocyclic aromatic groups such as phenylene, biphenylene, terphenylene, or naphthylene. Ar and Ar' may be the same or different.

Y is a substituent selected from organic, inorganic or organometallic radicals. The substituents represented by Y include (1) halogen, e.g., chlorine, bromine, iodine, or fluorine or (2) hydroxyl or ether groups of the general formula OE, wherein E is a monovalent hydrocarbon radical as defined below for X or (3) monovalent hydrocarbon groups as defined below for X or (4) other substituents, e.g., nitro or cyano, said substituents being essentially inert provided there be at least one and preferably two halogen atoms per arril e.g. phenyl nucleus

least one and preferably two halogen atoms per aryl, e.g., phenyl, nucleus.

X is a monovalent hydrocarbon group exemplified by the following: alkyl, such as methyl, ethyl, propyl, isopropyl, butyl, or decyl; aryl groups, such as phenyl, naphthyl, biphenyl, xylyl, or tolyl; aralkyl groups, such as benzyl, or ethylphenyl; cycloaliphatic groups, such as cyclopentyl, or cyclohexyl; as well as monovalent hydrocarbon groups containing inert substitutents therein. It will be understood that where more than one X is used they may be alike or different.

The letter d represents a whole number ranging from 1 to a maximum equivalent to the number of replaceable hydrogens substituted on the aromatic rings comprising Ar or Ar'. The letter e represents 0 or a whole number ranging from 1 to a maximum controlled by the number of replaceable hydrogens on R. The letters a, b, and c represent 0 or whole numbers. When b is not 0, neither a nor c may be 0. Otherwise either a or c, but not both, may be 0. Where b is 0, the aromatic groups are joined by a direct carbon-carbon bond

aromatic groups are joined by a direct carbon-carbon bond.

The Y substituents on the aromatic groups, Ar and Ar' can be varied in the ortho, meta or para positions on the aromatic rings and the groups can be in any

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	possible geometric relationship with respect to one another. Included within the scope of the above formula are di-aromatics of which the following are representative:	<u> </u>
_	2,2-bis-(3,5-dichlorophenyl)propane	
5	bis-(2-chlorophenyl)methane bis-(2,6-dibromophenyl)methane	5
	¹ , ¹ -DIS-(4-IOGODhenyl)ethane	·
	1,2-bis-(2,6-dichlorophenyl)ethane	
10	1,1-bis-(2-chloro-4-iodophenyl)ethane 1,1-bis-(2-chloro-4-methylphenyl)ethane	
	1,1°US-(3,3-CICRIOFORNERVI)ethane	10
	4,4-015-(3-DRCDVI-4-bromonhenyl)ethone	
	2,3-bis-(4,6-dichloronaphthyl)propane 2,2-bis-(2,6-dichlorophenyl)pentane	
15	4,4-018-(3,3-dichlorophenyl)hexane	
	018-(4-Chlorophenyl)phenylmethane	15
	bis-(3,5-dichlorophenyl)cyclohexylmethane bis-(3-nitro-4-bromophenyl)methane	
	UIST4-flyQfOXV-2.h-dichloro-3-mathovhht	
20	2,2-bis-(3,5-dichloro-4-hydroxyphenyl)propane 2,2-bis-(3-bromo-4-hydroxyphenyl)propane.	20
		20
	The preparation of these and other applicable biphenyls are known in the art. In place of the divalent aliphatic group in the above are related to the divalent aliphatic group	
25	sulfide, or sulfoxy.	
23	Included within the above structural con-	25
	biphenyls such as 2 2'-dichlorohiphenyl 3 4 111	25
	biphenyl, hexabromobiphenyl, october, 2,4-dioromobiphenyl, 2,4'-dichloro-	
30	halogenated diphenyl ethers, containing 2 to 10 halogen atoms. The preferred halogen companied for the preferred halogen atoms.	
	accordance with this invention only for use as flame-retardant additives in	30
	chlorinated benzene, brominated benzene, chlorinated biphenyl, chlorinated terphenyl, brominated biphenyl, brominated biphenyl, brominated biphenyl, brominated biphenyl, brominated terphenyl, brominated biphenyl, brominated terphenyl, brominated biphenyl, brominated biphenyl, brominated terphenyl, brominated biphenyl, brom	
	two phenyl radicals separated by a district to phenyl or a compound comprising	
35	chlorine or bromine atoms per phenyl nucleus, and mixtures of at least two foregoing.	
	Especially preferred as bout	35
	Especially preferred are hexabromobenzene and chlorinated biphenyls, alone, or mixed with antimony trioxide.	
0	in general, the preferred phosphoto	
	phosphonites phosphinites phosphinates	40
	phosphates Illustrative and the Control of the phosphines, phosphites or	
	antimony trioxide and optionally	
5	lypical of the preferred wheelers	
	invention would be those having the general formula	45
	o	
	QO — P — OQ OQ	
	Q0 — F — OQ	
	where each Q represents the same or different radicals including hydrocarbon	
0	alkyl: halogen, hydrogen and anyl substituted aryl and aryl substituted	
	said radicals O is any Typical are the feet provided that at least one of	50
	bisdodecyl phosphate, phenylbisneopentyl phosphate, phenylethyl hydrogen phosphate	

said radicals Q is aryl. Typical examples of suitable phosphates include, phenyl-bisdodecyl phosphate, phenylbisneopentyl phosphate, phenylethyl hydrogen phosphate, phenyl-bis-(3,5,5'-trimethylhexyl) phosphate, ethyldiphenyl phosphate, 2-ethylhexyl di(p-tolyl) phosphate, diphenyl hydrogen phosphate, bis(2-ethylhexyl) phosphate, tritolyl phosphate, bis(2-ethylhexyl) phenyl) phosphate, phenylmethyl hydrogen phosphate, di(dodecyl) p-tolyl phosphate, tricresyl phosphate, triphenyl phosphate, halogenated triphenyl

		9
5	phosphate, dibutylphenyl phosphate, 2-chloro-ethyldiphenyl phosphate, p-tolyl bis(2,5,5'-trimethylhexyl)phosphate, 2-ethylhexyldiphenyl phosphate, and diphenyl hydrogen phosphate. The preferred phosphates are those where each radical Q is aryl. The most preferred phosphate is triphenyl phosphate. It is also preferred to use triphenyl phosphate in combination with hexabromobenzene and, optionally, antimony trioxide. Also suitable as flame-retardant additions for an allowed the suitable and the suitable as flame-retardant additions for an allowed the suitable as flame-retardant additions for an allowed the suitable as flame-retardant additions for a suitable as flame-retardant additions for all suitable	5
10	Also suitable as flame-retardant additives for use in accordance with this invention are compounds containing phosphorus-nitrogen bonds, such as phosphonitrilic chloride, phosphorus ester amides, phosphoric acid amides, phosphonic acid amides, phosphonic acid amides, or tetrakis(hydroxymethyl)phosphonium chloride. These flame-retardant additives are commercially available.	10
15	The polytetrafluoroethylene resins used in accordance with this invention as dripping retarding agents (e) are commercially available or can be prepared by known processes. They are white solids obtained by free radical initiated polymerization of tetrafluoroethylene in aqueous media with free radical catalysts, e.g., sodium, potassium or ammonium peroxydisulfates at 100 to 1000 psi. at 0.200°C. and preferably at 20—100°C.	15
20	While not essential, it is preferred to use the resins in the form of relatively large particles, e.g., of average size 0.3 to 0.7 mm., mostly 0.5 mm. These are better than millimicrops in diameter. It is	20
25	them together into fibrous networks. Such preferred polytetrafluoroethylenes are designated by ASTM as Type 3, and are available commercially for general use in the extrusion of thin-walled tubular goods and tape.	25
30	usually will be from 0.1 to 10 parts and preferably from 0.5 to 2.5 parts by weight per hundred parts by weight of the combination of components (a) + (b). The reinforcements are added in any usual manner, e.g., by dry mixing or by mixing in the melted state in an extruder.	30
35	chopped into small pieces, e.g., 1" to 1" in length and put into an extrusion the flame retardant additive (d) and polyterafluoroethylene (e) to produce	35
40	optionally, flame retardant additive and polytetrafluoroethylene resin, by dry chopped. In still another procedure continuous lengths of glass roving are drawn	40
45	optionally, the flame retardant additive and polytetrafluoroethylene resin, which procedure coats the filaments, and then the resin-coated glass strand is mixed with resin and additives and directly molded, e.g., by injection or transfer	45
50	It is always very important to thoroughly free all of the ingredients, polyester resin, co-blending polymer, reinforcing filler, and flame retardant additives, from an addition, compounding should be as it is	50
55	heat is utilized; and an intimate blend between the resin and the additives is Although it is not essential best acres.	55
60	resin, the co-blending polymer and other additives, and the reinforcement, e.g., blend of the ingredients, the screw employed having a long transition section to	60
65	28 mm. Werner Pfleiderer machine can be fed with resins and additives at the feed port and reinforcement down stream. In either case, a generally suitable machine temperature will be 350 to 700°F.	
		65

	1.409.275	
	The precompounded composition can be extruded and cut up into pieces of	10_
5	The compositions can be molded in any equipment conventional	
10	of the Newbury type, with conventional cylinder temperatures, e.g., soo's, and conventional mold temperatures, e.g., 150°F. If necessary, depending filler and the recombination of the co-blending polymer, the amount of reinforcement	5
4.5	accommodate the composition. Typical conditions for a wide variety of materials The following accommodate the composition accommodate the composition.	10
15	The following examples illustrate the invention. They are set forth as a further description, but are not to be construed as limiting the invention thereto.	15
20	The following ingredients are dried in a vacuum at 100°C. for 10 hours: poly(1,4-butylene terephthalate), m.p., 232°C., intrinsic viscosity: 0.65—0.75	
20	ibrous glass reinforcement, 1", The dry blends are extruded and moulded between 470 and 520°F into test	20
25	The test bars are tested for the following physical properties: Tensue Strength and elongation, ASTM D-638; Flexural strength and modulus, ASTM D-790; The formulations used and the results obtained are set out in Table 1:	25

Table 1.

Physical Properties of Reinforced Polyester-Poly(methyl methacrylate)

Compositions

Example	ositions			
Ingredients	1A	ı	2	3
poly(1,4-butylene terephthalate)	0	20	40	60
poly(methyl methacrylate)	80	60	40	20
f fibrous glass reinforcement Properties	20	20	20	20
Heat Distortion temp., °F. at 264 psi	194	192	320	311
Flexural modulus, psi	660,000	811,000	851,000	750,000
Impact Strength, ft.lbs./in. notch	1.1	1.1	1.2	1.3

Thus, a composition comprising two polymers and a glass reinforcement is seen to be highly useful. Blends containing equal amounts of polyester and polymethacrylate show a surprising increase in heat distortion temperature, a maintenance of good impact strength and a very high flexural modulus.

30

The following ingredients are dried:
poly(1,4-butylene terephthalate) as in Example 1;
polyethylene, high density, 0.940—0.965 g/cc., (containing about 4° of fibrous glass reinforcement. 4"

EXAMPLES 4—6.

poly(1,4-butylene terephthalate) as in Example 1;
polyethylene, high density, 0.940—0.965 g/cc., (containing about 4° of fibrous glass reinforcement. 4"

10

The dry blends are extruded and molded at 520°F, into test pieces and tested for physical properties.

The formulations used and the results obtained are set out in Table 2:

Table 2. Physical Properties of Reinforced Polyester-Polyethylene Compositions

Example				
Ingredients (parts by weight)	4 A	4	•	
poly(1,4-butylene terephthalate)	 -		5	6
polyethylene	0	20	40	60
-	80	60	40	20
4" fibrous glass reinforcement	20	•		20
Properties	20	20	20	20
Heat distortion temp., °F. at 264 p	ei ara			
Flexural modulus, psi	SI. 212	235	262	392
•	273,000	275,000	311,000	348,000
Impact strength, notched, ftlbs/i	nch 2.7	1.7	1.5	1.2

5 When the formulation comprises equimolar amounts of polyester and polyethylene, a high heat distortion temperature and good notched impact strength were obtained. Also, as the amount of polyester is increased, both the modulus and

10 The following ingredients are dried: poly(1,4-butylene terephthalate), as in Example 1; acetal copolymer, density, 1.40 g/cc., (Celcon, manufactured by the Celanese EXAMPLE 7. Company); and 15

fibrous glass reinforcement, as in Example 1.

The dry blend is melt blended on a two-roll mill and compression molded at 220—230°C. The formulations used and the results obtained are set out in Table 3: 15

Table 3. Physical Properties of Reinforced Polyester-Acetal Copolymer Compositions

Example		
Ingradient	_7A	7
Ingredients (parts by weight)		
poly(1,4-butylene terephthalate)	0	40
acetal copolymer	_	40
fibrous glass reinforcement	80	40
Properties	20	20
Heat Distortion temp., °F. at 264 psi.	20.	
Modulus, psi.	304	313
чаская, раг.	600,000	570,000

10

The composite according to this invention has an increased heat distortion temperature in comparison with the reinforced acetal copolymer alone.

EXAMPLES 8 to 13.

The following ingredients are dried: The following ingredients are dried:
poly(1,4-butylene terephthalate), as in Example 1;
polyamide resin, nylon 6/10, density, 1.09 g/cc. (Zytel 31, manufactured by
DuPont Company, "Zytel" is a Registered Trade Mark);
polyamide resin, nylon 6, density, 1.12—1.14 g/cc.; and
fibrous glass reinforcement, as in Example 1.
The dry blends of nylon 6/10 are melt blended on a two-roll mill at
210—220°C. and compression molded at 230°C. The dry blends of nylon 6 are
extrusion blended at 520°F. The formulations used and the results obtained are set
out in Table 4: 5 10

Table 4. Physical Properties of Reinforced Polyester-Polyamide Compositions

Example	8A	8	. 9	10
Ingredients (parts by weight)				
poly(1,4-butylene terephthalate)	0	20	40	60
polyamide, nylon 6	80	60	40	20
f fibrous glass reinforcement	20	20	20	20
Properties				0
Heat distortion temp., ° F. at 264 psi	379	383	410	406
Flexural strength, psi.	6,700	8,400	7,100	5,700
Modulus, psi.	178,000	330,000	500,000	530,000
Impact strength, ftlbs./in. notch	3.6	2.0	1.5	1.3

¹⁵ As nylon 6 is blended into the polyester, easier processing is obtained. At the 60:20 ratio of nylon to polyester, the modulus increases sharply. This is also true of the 40-40 blend.

15

10

5

10

Table 4 (continued). Physical Properties of Reinforced Polyester-Polyamide Compositions

Example	11A	11		
Ingredients (parts by weight)				
poly(1,4-butylene terephthalate	0	20	40	60
polyamide, nylon 6/10	80	20	40	60 20
f" fibrous glass reinforcement	20		20	20
Properties			20	20
Heat distortion temp., °F. at 264 psi	408	414	400	421
Flexural strength, psi.	14,000	14,300	6,000	5,200
Modulus, psi.	400,000	447,000	455,000	356,000
Impact strength, ftlbs./in. notch	1.2	1.4	1.4	2.3

Improved processing is obtained at the 60-20 blend of polyester and polyamide with increased impact strength and heat distortion temperature.

EXAMPLES 14 to 16.

EXAMPLES 14 to 16.

The following ingredients are dried:
poly(1,4-butylene terephthalate), as in Example 1;
composition comprising poly(2,6-dimethyl-1,4-phenylene) ether and rubber modified, high impact polystyrene, equal parts density, 1.05—1.10; and fibrous glass reinforcement, ‡ inch, (P 158 B, manufactured by Owens Corning Fiberglas Corp.).

The dry blends are melt blended by extrusion at 460—520°F. and molded at 520°F. The formulations used and the results obtained are set out in Table 5:

Table 5. Physical Properties of Reinforced Polyester-Polyphenylene Ether Compositions

Examples	14	16	
Ingredients (parts by weight)	14	15	. 16
poly(1,4-butylene terephthalate)	20	40	60
polyphenylene ether	30	20	10
polystyrene (rubber modified)	30	20	10
fibrous glass reinforcement	20	20	20
Properties			20
Heat distortion temp., °F. at 264 psi	260	298	395
Tensile strength, psi.	6,040	6,400	7,500
Flexural modulus, psi.	474,000	580,000	620,000
Impact strength ftlbs./inch, notch	1.1	1,2	1.7

	1,409,275	1.4
5	Improved extrusion is noticed as the amount of polyester is increased. In addition the composities according to this invention have a smoother molded surface than those containing no polyester. Good impact strengths are obtained with 10:10:60 polystyrene-polyphenylene ether-polyester combinations along with very high heat distortion temperature and flexural modulus. The procedure is repeated, substituting for the polyphenylene ether-styrene resin composition, an unmodified poly(2,6-dimethyl-1,4-phenylene)ether, (PPO, manufactured by General Electric Co.). Compositions according to this invention are obtained.	145
10	EXAMPLE 17. A dry blend of 60 parts by weight of poly(1,4-butylene terephthalate), 20 parts by weight of an (85-15 p-phenylene isophthalate terephthalate) wholly aromatic polyester, prepared according to U.S. Patent No. 3,036,990, and 20 parts by weight of fibrous glass reinforcement, as in Example 1 is milled at 240°C. and heated at 300°C. provide a moldable reinforced composition according to this invention.	10
15	EXAMPLES 18 to 22. The following ingredients are dried: poly(1,4-butylene terephthalate), as in Example 1; polyethylene as in Example 4.	15
20	highly chlorinated biphenyl (chlorine content 59%, softening point 98—105°C); hexabromobenzene; triphenyl phosphate; antimony trioxide: and	20
25	polytetrafluoroethylene resin (ASTM Type 3, particle size 0.3—0.7 mm.) The blends are compounded and molded according to the procedure of Example 1. The formulations used are set out in Table 6:	25

Table 6. Flame Retardant Reinforced Polyester-Polyethylene Compositions

Examples			
Ingredients (parts by weight)	18	19	20
- ·			
poly(1,4-butylene terephthalate)	20	20	20
polyethylene	60	60	60
fibrous glass reinforcement	20	20	20
chlorinated biphenyl	10	_	_
hexabromobenzene	_	7	
triphenyl phosphate		,	_
antimony trioxide	_	_	7
polytetrafluoroethylene resin	4	3	
- Joseph Todan	1.0	1.5	1.0

Table 6.

Flame Retardant Reinforced Polyester-Polyethylene
Compositions (continued)

Examples	21	
Ingredients (parts by weight)	21	22
poly(1,4-butylene terephthalate)	20	20
polyethylene	60	60
fibrous glass reinforcement	20	
chlorinated biphenyl hexabromobenzene	20	20
triphenyl phosphate		10.5
antimony trioxide	8	4.5
polytetrafluoroethylene resin	0.1	1.5

Flame retardant, non-dripping compositions according to this invention are obtained.

5	EXAMPLE 23. The procedure of Example 4 is repeated, substituting for the poly(1,4-able linear polyesters:	5
10	poly(1,3-propylene terephthalate-1,4-butylene isophthalate polyester; dimethyl terephthalate by the procedure of U.S. Patent No. 2,465,319, Example	10
15	poly(hexamethylene terephthalate) prepared from hexamethylene glycol and dimethyl terephthalate by the procedure of U.S. Patent No. 2,465,319, m.p., Reinforced compositions according to this invention are obtained.	
	For example, reinforced compositions according to this invention are obtained if, instead of polyethylampositions according to this invention are	15
20	Chemical Industries); a copolymer of ethylene and estylene and estylen	20
25	poly(vinyl chloride—vinyl acetate), density 1.35—1.45 g/cc.; poly(vinylidene chloride), density 1.65—1.72 g/cc.; natural rubber: and	25
30	a rubbery copolymer of butadiene (83%) and acrylonitrile (18%). In addition, there can be substituted for polyethylene in Example 4: a polyacetal homopolymer, density 1.42 g/cc.; a polysulfone, density 1.24 g/cc. (Polyella J. Cc.)	
	a polysulfone, density, 1.24 g/cc.; Carbide—Bakelite is a Registered Trade Mark); a polyamide; a polyorganosiloxane; copolymers of a polyorganosiloxane with, respectively, styrene, methyl	30
35	methacrylate and the reaction product of bisphenol-A and isophthaloyl chloride; a siloxane-nitrogen copolymer containing amido, amide-imido or imide groups.	35

	1,00,10	16
•	The procedure of Example 4 can be repeated, substituting for glass fibers, the following reinforcing fillers: aliminum powder;	
5	bronze powder; silicate;	
	ceramic fibers; titanate fibers; fumed colloidal silica;	5
10	asbestos fibers;	
10	quartz; and carbon black.	10
	Reinforced composites according to this invention are about the	
15	thermal properties and the enhanced flame resistance of certain embodiments, the reinforced polyester-coblending polymer composites of this invention have many and varied uses. The formulations may be used alone as molding powders or mixed with other polymers and may contain properties and may contain the second of the composition of	15
20	and plasticizers.	
20	WHAT WE CLAIM IS:—	20
	1. A reinforced thermoplastic composition that is rigid at temperatures of up to 90°F, which comprises a homogeneous crystalline normally rigid, reinforceable mixture of:	20
25	(a) at least one high molecular weight normally crystalline polyester resin which rapidly crystallizes from the melt (as hereinbefore defined); (b) at least one high molecular weight normally crystalline, normally amorphous or normally partially crystalline and partially amorphous polymer of: (i) at least one aliphatic ethylenically unsaturated monomer excluding propylene:	25
30	(ii) at least one difunctionally reactive compound which is polymerizable by self-condensation: and/or	
		30
	(iii) at least two difunctionally reactive compounds which are polymerizable by polycondensation;	
35	in the concentration range of from 1 to 99 parts by weight of (a) to 99 to 1 parts by weight of (b); and	
	(c) from 2 to 90% by weight, based on total composition, of a reinforcing filler for said homogeneous, crystalline, normally rigid, reinforceable mixture; and when plolymer (b) is a polyester, it is a wholly aromatic polyester and different from polyester (a).	35
40	2. A composition as defifned in Claim I wherein said polyester resin (a) is a poly(alkylene terephthalate, isophthalate or mixed terephthalate and isophthalate), said alkylene groups containing from 3 to 10 carbon atoms. 3. A composition as defined in Claim I wherein polyester resin (a) includes repeating units of the formula:	40
45	-0 - (CH ₂)4-0- C	45
	4. A composition as defined in Claim 3 wherein said polyester (a) is poly(1,4-butylene terephthalate).	
	5. A composition as claimed in any preceding claim wherein said polymer (b) is a polymerization product of at least one clieb at the said polymer (b)	
50	monomer and is polyethylene.	50
	poly(methyl methacrylate) poly(methylpentene),	
55	a copolymer of ethylene and ethyl acrylate, poly(vinyl chloride),	
	poly(vinylidene chloride)	55
	a copolymer of vinyl chloride and vinyl acetate, natural rubber, or a rubbery copolymer of butadiene and acrylonitrile.	

		17
	6. A composition as defined in any of Claims 1 to 4 wherein said polymer (b) is a polyacetal,	
	an acetal copolymer, a polyphenylene ether,	
5	a polysulfone,	
	a polyamide.	5
	a polyimide,	
	a polyorganosiloxane,	
10	a copolymer of a polyorganosiloxane and a vinyl aromatic monomer, an acrylic monomer or an aromatic ester, or	
10	a siloxane-nitrogen copolymer containing amido, amide-imido or imide groups.	10
	groups. amide-imido or imide	
	7. A composition as defined in any of claims 1 to 4 wherein said ethylenically unsaturated monomer is styrene and said diffuscionally assets.	
15	unsaturated monomer is styrene and said difunctionally reactive compound is 2,6-	
. 13	8. A composition as defined in our second in the second in	15
	8. A composition as defined in Claim 1 wherein said reinforcing filler (c) is a reinforcing metal, ceramic, silica, quartz, glass or carbon. 9. A composition as defined in Cl., glass or carbon.	
	9. A composition as defined in Claim a vi caroll.	
20	filamentous glass.	
20	10. A composition as defined in any preceding claim wherein said polyester resin is normally flammable, and the composition also include:	20
	(d) a flame retardant addition in composition also includes	24
	sufficient to render said polyester resin non-burning or self-extinguishing.	
	11. A composition as defined in Claim 10 which comprises: (e) a polytetrafluoroethylene regin in the comprises:	
25	(e) a polytetrafluoroethylene resin in a minor proportion based on said composition but in an amount at least sufficient to render raid not necessarily as the said ne	25
	dripping, when hurning	43
	12. A composition as defined in Claim to	
	additive is a halogen-containing compound; a halogen-containing compound in	
30	admixture with antimony trioxide; elemental phosphorus, an organic phosphonic	30
	acid, a phosphonate, phosphinate, a phosphonic, a phosphinite, a phosphine, a phosphine, a phosphine, a phosphine oxide, a phosphine, a phosphite, a phosphine or a nature of any of the foregoing.	30
	13. A composition as defined in Clarific of a maxture of any of the foregoing.	
	additive is triphenyl phosphate.	
35	14. A composition as defined in Claim 12 wherein said flame retardant additive is chlorinated benzene brominated benzene	25
	chlorinated tempenyl brominated benzene, chlorinated biphenyl.	35
	comprising two phenyl radicals compound	
	and having at least two chlorine or bromine substituents per benzene ring, or a mixture thereof.	
40	15. A composition of definition of the state	40
	15. A composition as defined in Claim I substantially as hereinbefore described.	40
	• • • • • •	

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